(1) Publication number:

0 020 037

B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication of patent specification: 15.08.84

(1) Application number: 80301487.7

② Date of filing: 07.05.80

(5) Int. Cl.³: C 10 M 1/36, C 10 M 3/30,

C 10 L 1/22,

C 07 C 103/14,

C 07 D 207/40

(54) Oil-soluble friction-reducing additive, process for the preparation thereof, and lubricating oil or fuel composition containing the additive.

- (3) Priority: 18.05.79 US 40413
- (43) Date of publication of application: 10.12.80 Bulletin 80/25
- (45) Publication of the grant of the patent: 15.08.84 Bulletin 84/33
- (84) Designated Contracting States: BE DE FR GB IT NL
- (58) References cited: GB - A - 877 845 GB - A - 1 111 837 US - A - 3 382 172 US - A - 3 655 351

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Description

This invention is in the field of lubricants and lubricant additives. More particularly, the invention deals with additives to lubricants especially for crankcase use for internal combustion engines, which provide a reduction of friction of the operating engine.

In order to conserve energy, automobiles are now being engineered to give improved gasoline mileage compared to those in recent years. This effort is of great urgency in the United States in view of regulations which compel auto manufacturers to achieve prescribed gasoline mileage. These regulations are to conserve crude oil. In an effort to achieve the required mileage, new cars are being downsized and made much lighter. However, there are limits in this approach beyond which the cars will not accommodate a typical family.

Another way to improve fuel mileage is to reduce engine friction. The present invention is concerned with this latter approach.

The applicants draw attention to U.S. Patent Specification No. 3382172 and to British Patent Specification No. 1111837. The British Specification refers to C_{8-42} alkyl-substituted cyclic imides, including succinimides, which may be used as lubricating oil additives. It is however, pointed out that this British Patent Specification is not concerned with cyclic imides in which the substituents are unsaturated and joined to the ring by a secondary attachment nor with compounds made using the isomerisation technique which is essential in the present invention. U.S. Patent Specification No. 3382172 refers to C_{14-22} alkenyl succinic acids which may be used as antiwear lubricating oil additives. This U.S. Specification does not, however, disclose imide or amide products nor products having a friction reducing effect.

The present invention utilizes the finding that oil soluble, C₁₂₋₃₆ aliphatic hydrocarbyl succinimides or succinimides containing a certain hydrocarbon grouping which has been formed by a particular isomerization technique provide a friction reducing effect when they are incorporated in lubricating oil.

Accordingly, the present invention provides an oil-soluble friction-reducing additive which comprises at least one compound having the structure:

$$Z - CH - C$$
 $Z - CH - C - NH_2$
 $CH_2 - C - NH_2$

or
$$H_2N$$
 C CHZ CH_2 CH_2 CH_3 C CH

wherein

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n is an integer of from 2 to 4 and wherein Z has the structure:

wherein

 R_1 and R_2 are each independently straight or branched chain hydrocarbon groups containing from 1 to 34 carbon atoms such that the total number of carbon atoms in the groups R_1 and R_2 is from 11 to 35, the Z hydrocarbon group being unsaturated and formed by reacting maleic acid, anhydride or ester with an internal olefin containing 12 to 36 carbon atoms, said internal olefin being formed by isomerizing the olefinic double bond of a linear α -olefin or mixture thereof to obtain a mixture of internal olefins.

The aliphatic hydrocarbon group is derived from a linear α -olefin which has been isomerized to form a mixture of internal olefins. The additive can also be used in the engine fuel.

A preferred embodiment of the invention is a lubricating oil composition or fuel composition containing a friction-reducing amount of an additive as above defined.

The aliphatic substituent Z on the succinic group can be any unsaturated aliphatic hydrocarbon group containing from 12 to 36 carbon atoms including alkenyl and polyunsaturated hydrocarbon groups.

Examples of the above additive include:

1,2-dimethyl octadecenyl succinimide

1-methyl-3-ethyl dodecenyl succinimide

In a highly preferred embodiment R_1 and R_2 are straight chain aliphatic hydrocarbon groups. These additives have improved solubility in lubricating oil. Examples of these additives are:

1-propyltridecenyl succinimide

1-pentyltridecenyl succinimide

1-tridecylpentadecenyl succinimide

1-tetradecyleicosenyl succinimide

The above highly preferred additives are made from linear α -olefins containing from 12 to 36 carbon atoms by isomerizing the α -olefins to form a mixture of internal olefins and reacting this mixture of internal olefins with maleic acid, anhydride or ester forming an intermediate and reacting the intermediate with ammonia to form amide, imide, or mixtures thereof.

Additives made from isomerized linear α -olefins have greatly improved oil solubility compared with additives made with linear α -olefins.

Isomerization of the linear α -olefin can be carried out using conventional methods. One suitable method is to heat the linear α -olefin with an acidic catalyst. Especially useful acid catalysts are the sulfonated styrenedivinylbenzene copolymers. Such catalysts are commercially available and are conventionally used as cation exchange resins. In the present method they are used in their acid form. Typical resins are Amberlyst 15, XN—1005 and XN—1010 (registered trademarks) available from Rohm and Haas Company. Use of such resins for isomerizing linear α -olefins is described in US—A—4,108,889.

The method by which the present additives are made are illustrated by the following examples.

Example 1

In a reaction vessel was placed 1000 grams of linear α -octadecene. To this was added 187 grams Amberlyst 15 (5 percent moisture). The mixture was stirred under nitrogen and heated at 120°C for 3 hours. The isomerized product contained 3.6 weight percent olefin dimer and the balance was internal C_{18} olefin. The product was separated from the resin.

In a second reaction vessel was placed 504 grams of the above isomerized C₁₈ olefin and 300 ml heptane. The heptane was distilled out under vacuum to remove water. Then 2.4 grams of tri-(3,5-di-tert-butyl-4-hydroxybenzyl)-mesitylene stabilizer was added. The mixture was heated under nitrogen to 225°C. Then 160 grams of molten maleic anhydride was slowly added over a 2.5 hour period. The mixture was stirred at 225°C for two more hours and then unreacted maleic anhydride was distilled out by pulling vacuum to 1.0132M Pa (762 mmHg) (30 in.) while holding the reaction mixture at 200°C. The product was principally secondary C. alkylene succinic aphydride.

The product was principally secondary C₁₈ alkylene succinic anhydride.

In a separate reaction vessel was placed 532.5 grams of the above isomerized octadecenyl succinic anhydride. This was heated under nitrogen to 165°C and then ammonia was injected causing the temperature to rise to 180°C. Ammonia injection was continued until exotherm stopped. The mixture was heated at 170°C under vacuum to remove water yielding isomerized octadecenyl succinimide.

Example 2

In a reaction vessel was placed 1005 grams of linear α -eicosene and 187 grams of Amberlyst 15 (5 percent moisture). The mixture was heated under nitrogen at 110° to 125°C for 6 hours. The product was internally unsaturated eicosene containing 3.3 percent eicosene dimer.

In a separate reaction vessel was placed 560 grams of the above isomerized eicosene and 200 ml heptane. The heptane was distilled out to dry the eicosene. At 140°C, 3.1 grams of tri-(3,5-di-tert-butyl-4-hydroxybenzyl)mesitylene stabilizer was added and the mixture heated to 210°C. Over a 2.5 hour period, 156.8 grams of maleic anhydride was added at about 225°C. Following this, unreacted maleic anhydride was distilled out under vacuum at 210°C. Leaving isomerized eicosenyl succinic anhydride.

In another reaction vessel was placed 570 grams of the above isomerized eicosenyl succinic anhydride. This was heated to 160°C and ammonia injection started. The temperature rose to 175°C. Ammonia injection was continued at 175°C until the temperature dropped. Then 1.0132M Pa (762 mm Hg) (30 in.) vacuum was slowly applied to distill out water and ammonia. Additional ammonia was injected to be sure no anhydride remained. There was no further reaction so this ammonia was stripped out at 1.0132M Pa (762 mm Hg) (30 in.) vacuum at 170°C yielding isomerized eicosenyl succinimide.

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Example 3

In a reaction vessel was placed 1100 grams of linear C_{16} — C_{18} α -olefin mixture. The olefin mixture was isomerized following the procedure in Example 2.

In a separate vessel was placed 485 grams (2 moles) of the above isomerized olefin. This was heated at 100°C under 1.0132M Pa (30 inches Hg) vacuum to remove water. To it was then suided 2.4 grams tri-(3,5-di-tert-butyl-4-hydroxybenzyl)mesitylene. The mixture was heated under birogen to 225°C and then 152 grams of molten maleic anhydride was added over a 3 hour period. The mixture was stirred 30 minutes at 225°C and an additional 50 grams of maleic anhydride was ended. This mixture was stirred 30 minutes at 225°C following which unreacted maleic anhydride was added out at 200°C under 1.0132M Pa (762 mm Hg) (30 in) vacuum.

In a separate reaction vessel was placed 598 grams of the above isomerized C_{19} — 1_{10} is identification was started at 140°C raising the temperature 1_{10} .45°C. Ammonia injection was continued at 130°C until no further ammonia was adsorbed. The resture was then heated to 180°C to distill out water and ammonia yielding isomerized C_{10} — C_{18} alkenyl succinimide.

The additives are added to the lubricating oil in an amount which reduces the fraction of the engine operating with the oil in the crankcase. A useful concentration is about 0.05 to 3 weight percent. A more preferred range is about 0.1 to 1.0 weight percent.

From the above it can be seen that the present invention provides an improved prankcase lubricating oil. Accordingly, an embodiment of the invention is an improved motor oil composition formulated for use as a crankcase lubricant in an internal combustion engine wherein the improvement comprises including in the crankcase oil an amount sufficient to reduce fuel consumption of the engine of the friction-reducing additive herein described.

In a highly preferred embodiment such improved motor oil also contains an ashless dispersant and an alkaline earth metal salt of a petroleum sulfonic acid or an alkaryl sulfonic acid (e.g. alkyibenzene sulfonic acid).

The additives can be used in mineral oil or in synthetic oils of viscosity suitable for use in the crankcase of an internal combustion engine. Crankcase lubricating oils have a viscosity up to about 16 δ 10⁻³ m²/S (80 SUS) at 100°C (210°F).

Mineral oils include those of suitable viscosity refined from crude oil from all sources including Gulfcoast, midcontinent, Pennsylvania, California, Alaska and the like. Various standard refinery operations can be used in processing the mineral oil.

Synthetic oil includes both hydrocarbon synthetic oil and synthetic esters. Useful synthetic hydrocarbon oils including liquid polymers of α -olefins having the proper viscosity. Especially useful are the hydrogenated liquid oligomers of C_6 — C_{12} α -olefins such as α -decene trimer. Likewise, alkylbenzenes of proper viscosity can be used, such as didoecylbenzene.

Useful synthetic esters include the esters of both monocarboxylic acid and polycarboxylic acid as well as monohydroxy alkanols and polyols. Typical examples are didoceyl adipate, trimethylol propane tripelargonate, pentaerythritol tetracaproate, di(2-ethylhexyl)adipate, dilauryl sebacate and the like. Complex esters prepared from mixtures of mono- and dicarboxylic acid and mono- and polynydroxyl alkanols can also be used.

Blends of mineral oil with synthetic oil are particularly useful. For example, blends of 5 to 25 weight percent hydrogenated α -decene trimer with 75 to 95 weight percent 32 x 10⁻³ m²/S (150 SUS 38°C (100°F)) mineral oil results in an excellent lubricant. Likewise, blends of about 5 to 25 weight percent di(2-ethylhexyl)adipate with mineral oil of proper viscosity results in a superior lubricating oil. Also blends of synthetic hydrocarbon oil with synthetic esters can be used. Blends of mineral oil with synthetic oil are especially useful when preparing low viscosity oil (e.g., SAE 5W 20) since they permit these low viscosities without contributing excessive volatility.

The more preferred lubricating oil compositions include zinc dihydrocarbyldithiophosphate (ZDDP) in combination with the present additives. Both zinc dialkyldithiophosphates and zinc dialkaryldithiophosphates as well as mixed alkyl-aryl dithiophosphates can be used. Examples of alkyl-type ZDDP are those in which the hydrocarbyl groups are a mixture of isobutyl and isoamyl alkyl groups. Zinc di-(nonylphenyl)-dithiophosphate is an example of an aryl-type ZDDP. Good results are achieved using sufficient zinc dihydrocarbyldithiophosphate to provide about 0.01 to 0.5 weight percent zinc. A preferred concentration supplies about 0.05 to 0.3 weight percent zinc.

Another additive used in the oil compositions are the alkaline earth metal petroleum sulfonate or alkaline earth metal alkaryl sulfonates. Examples of these are calcium petroleum sulfonates, magnesium petroleum sulfonates, barium alkaryl sulfonates, calcium alkaryl sulfonates or magnesium alkaryl sulfonates. Both the neutral and the overbased sulfonates having base numbers up to about 400 can be beneficially used. These are used in an amount to provide about 0.05 to 1.5 weight percent alkaline earth metal and more preferably about 0.1 to 1.0 weight percent.

Viscosity index improvers can be included such as the polyalkylmethacrylate type or the ethylene-propylene copolymer type. Likewise, styrene-diene VI improvers can be used. Alkaline earth metal salts of phosphosulfurized polyisobutylene are useful. Preferred crankcase oils also contain an ashless dispersant such as the polyolefin succinamides and succinimides of polyethylene polyamines such as

tetraethylenepentamine. The polyolefin succinic substituent is preferably a polyisobutene group having a molecular weight of from about 800 to 5,000. Such ashless dispersants are more fully described in U.S. 3,172,892 and U.S. 3,219,666 incorporated herein by reference.

Other useful ashless dispersants include the Mannich condensation products of polyolefin-substituted phenols, formaldehyde and polyethylene polyamine. Preferably, the polyolefin phenol is a polyisobutylene-substituted phenol in which the polyisobutylene group has a molecular weight of from about 800 to 5,000. The preferred polyethylene polyamine is tetrathylene pentamine. Such Mannich ashless dispersants are more fully described in U.S. 3,368,972; U.S. 3,413,347; U.S. 3,442,808; U.S. 3,448,047; U.S. 3,539,633; U.S. 3,591,598; U.S. 3,600,372; U.S. 3,634,515; U.S. 3,697,574; U.S. 3,703,536; U.S. 3,704,308; U.S. 3,725,480; U.S. 3,726,882; U.S. 3,736,357; U.S. 3,751,365; U.S. 3,756,953; U.S. 3,793,202, U.S. 3,798,165; U.S. 3,798,247 and U.S. 3,803,039.

The friction-reducing additives of this invention are also useful in fuel compositions. Fuel injected or inducted into a combustion chamber wets the walls of the cylinder. Fuels containing a small amount of the present additive reduce the friction due to the piston rings sliding against the cylinder wall.

The additives can be used in both diesel fuel and gasoline used to operate internal combustion engines. Fuels containing about 0.001 to 0.25 weight percent of the friction-reducing additives can be used.

Fuels used with the invention can contain any of the additives conventionally added to such fuels. In the case of gasoline it can include dyes, antioxidants, detergents, antiknocks (e.g. tetraethyllead, methylcyclopentadienylmanganese tricarbonyl, rare earth metal chelates, methyl *tert*-butylether and the like). In the case of diesel fuels the compositions can include pour point depressants, detergents, ignition improvers (e.g. hexanitrate) and the like.

Tests were conducted using a 1977 U.S. production automobile. These were shortened versions of the Federal City EPA cycle. This is referred to as the "Hot 505" cycle. It consists of the first 5.8 km (3.6 miles) of the Federal EPA City cycle started with a warmed-up engine instead of a cold engine. The car with a fully formulated SE grade oil in its crankcase is operated on a chassis dynamometer for about one hour at about 88 km/hr. (55 mph) to stabilize oil temperature. It is then run through four consecutive "Hot 505" cycles measuring fuel economy of the base oil. Results of the four cycles are averaged. Then one-half of the base oil is drained from the crankcase and replaced with the same base oil containing a double dose of the test additive. The car is then run at about 88 km/hr. (55 mph) for about one hour to again stabilize temperature. A second series of four consecutive "Hot 505" cycles is run to measure initial fuel economy of the base oil containing the test additive. The car is then run about 805 km (500 miles) at constant speed of about 88 km/hr. (55 mph). Then a third series of four consecutive "Hot 505" cycles are run to measure fuel economy after about 805 km (500 miles) operation on the oil containing the test additive. The crankcase is then drained hot and filled with flushing oil which is run for a short time and then drained. The crankcase is then filled with the base oil which is run for a short time and then drained. The crankcase is then filled a second time with a base oil. This is run about one hour at about 88 km/hr. (55 mph) to a stable temperature. Then a fourth series of four consecutive "Hot 505" cycles are run measuring fuel economy. This gives a second base line thus bracketing the test carried out with the friction additive between two base line tests.

The following table shows the percent improvement in fuel economy over the base oil obtained using 1 weight percent of the friction-reducing additive.

| Additive | Percent Gain in Fuel Economy | | |
|---|------------------------------|--------------------------|---------|
| | Initial | About 805 km (500 miles) | Average |
| Isomerized C ₁₆₋₁₈ alkenyl succinimide | 1.6 | 0.9 | 1.3 |

The reduction in fuel consumption though small is significant.

Claims

1. An oil-soluble friction-reducing additive which comprises at least one compound having the structure:

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$$z - CH - C$$
 $CH_2 - C$
 $CH_2 -$

or:
$$H_2N = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} = CHZ = CH_2 = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} = NH = H$$

wherein

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n is an integer of from 2 to 4 and wherein Z has the structure:

wherein

 R_1 and R_2 are each independently straight or branched chain hydrocarbon groups containing from 1 to 34 carbon atoms such that the total number of carbon atoms in the groups R_1 and R_2 is from 11 to 35, the Z hydrocarbon group being unsaturated and formed by reacting maleic acid, anhydride or ester with an internal olefin containing 12 to 36 carbon atoms, said internal olefin being formed by isomerizing the olefinic double bond of a linear α -olefin or mixture there to obtain a mixture of internal olefins.

2. An oil-soluble friction-reducing additive as claimed in claim 1 wherein $\rm R_1$ and $\rm R_2$ are straight chain aliphatic hydrocarbon groups having a total of from 15 to 21 carbon atoms.

3. An oil-soluble friction reducing additive as claimed in claim 1 which is 1-propyltridecenyl succinimide, 1-pentyltridecenyl succinimide, 1-tridecylpentadecenyl succinimide or 1-tetradecyleicosenyl succinimide,

4. A process for the preparation of an oil-soluble additive as claimed in claim 1 which process comprises (a) isomerizing the olefinic double bond of a linear α -olefin or a mixture thereof containing from 12 to 36 carbon atoms to obtain a mixture of internal olefins, (b) reacting the mixture of internal olefins with maleic acid, anhydride or ester to obtain an intermediate hydrocarbon-substituted succinic acid, anhydride or ester and (c) reacting the intermediate with ammonia to form an amide, imide or a mixture thereof.

5. A process as claimed in claim 4 wherein the α -olefin consists mainly of linear α -tetradecene, linear α -hexadecene, linear α -octadecene or linear α -eicosene.

6. A process as claimed in claim 4 or claim 5 wherein the isomerization of the linear α -olefin is carried out by heating the linear α -olefin with an acidic catalyst.

7. A lubricating oil or fuel composition which contains a friction-reducing amount of an additive as claimed in any one of claims 1 to 3 or whenever prepared by a process as claimed in any one of claims 4 to 6.

8. A lubricating oil or fuel composition as claimed in claim 7 wherein the additive is tetradecenyl succinimide, hexadecenyl succinimide, octadecenyl succinimide or eicosenyl succinimide.

 A lubricating oil composition as claimed in claim 7 or claim 8 wherein the additive is contained therein in an amount of from 0.05 to 3 weight percent.

10. A fuel composition as claimed in claim 7 or claim 8 wherein the additive is contained therein in an amount of from 0.001 to 0.25 weight percent.

Patentansprüche

 Öllösliches reibungsminderndes Zusatzmittel, welches mindestens eine Verbindung der folgenden Strukturen enthält

oder
$$H_2N = \begin{bmatrix} O \\ \parallel \\ C - CHZ - CH_2 - C - NH \end{bmatrix}$$

worin n eine ganze Zahl von 2 bis 4 ist und worin Z die Struktur

besitzt, worin R_1 und R_2 jeweils unabhängig geradkettige oder verzweigtkettige Kohlenwasserstoffreste mit 1 bis 34 Kohlenstoffatomen derart sind, daß die Gesamtzahl der Kohlenstoffatome in den Resten R_1 und R_2 von 11 bis 35 beträgt, der Kohlenwasserstoffrest Z ungesättigt ist und durch Umsetzung von Maleinsäure, -anhydrid oder -ester mit einem inneren Olefin, welches 12 bis 36 Kohlenstoffatome enthält, hergestellt wird, wobei das innere Olefin durch Isomerisierung der olefinischen Doppelbindung eines linearen α -Olefins oder eines Gemisches davon gebildet wird, um ein Gemisch innerer Olefine zu erhalten.

2. Öllösliches reibungsminderndes Zusatzmittel nach Anspruch 1, worin R_1 und R_2 geradkettige aliphatische Kohlenwasserstoffreste mit einer Gesamtzahl von 15 bis 21 Kohlenstoffatomen sind.

3. Öllösliches reibungsminderndes Zusatzmittel nach Anspruch 1, welches 1-Propyltridecenylsuccinimid, 1-Pentyltridecenylsuccinimid, 1-Tridecylpentadecenylsuccinimid oder 1-Tetradecyleicosuccinimid ist,

4. Verfahren zur Herstellung eines öllöslichen Zusatzmittels nach Anspruch 1, wobei (a) die olefinische Doppelbindung eines linearen α -Olefins oder eines Gemisches davon mit 12 bis 36 Kohlenstoffatomen isomerisiert wird, um ein Gemisch von inneren Olefinen zu erhalten, (b) das Gemisch von inneren Olefinen mit Maleinsäure, -anhydrid oder -ester umgesetzt wird, um eine intermediäre Kohlenwasserstoff-substituierte Succinsäure, -anhydrid oder -ester zu erhalten und (c) die Zwischenverbindung mit Ammoniak umgesetzt wird, um ein Amid, Imid oder ein Gemisch davon zu bilden.

5. Verfahren nach Anspruch 4, worin das α -Olefin hauptsächlich aus linearem α -Tetradecen, linearem α -Hexadecen, linearem α -Octadecen oder linearem α -Eicosen besteht.

6. Verfahren nach Anspruch 4 oder Anspruch 5, worin die Isomerisierung des linearen α -Olefins durch Erhitzen des linearen α -Olefins mit einem sauren Katalysator durchgeführt wird.

7. Schmieröl-oder Brennstoffzusammensetzung, die eine reibungsmindernde Menge eines Zusatzmittels nach einem der Ansprüche 1 bis 3 enthalten oder nach einem Verfahren gemäß einem der Ansprüche 4 bis 6 hergestellt sind.

8. Schmieröl-oder Brennstoffzusammensetzung nach Anspruch 7, worin das Zusatzmittel Tetradecenylsuccinimid, Hexadecenylsuccinimid, Octadecenylsuccinimid oder Eicosenylsuccinimid ist.

9. Schmieröl-Zusammensetzung nach Anspruch 7 oder Anspruch 8, worin das Zusatzmittel darin in einer Menge von 0,05 bis 3 Gew.-% enthalten ist.

10. Brennstoff-Zusammensetzung nach Anspruch 7 oder Anspruch 8, worin das Zusatzmittel darin in einer Menge von 0,001 bis 0,25 Gew.-% enthalten ist.

Revendications

1. Additif soluble dans l'huile, réduisant la friction, qui comprend au moins un composé de formule:

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$$z - CH - C$$
 $CH_2 - CH_2$
 $CH_2 - CH_2$
 $CH_2 - CH_2$
 $CH_2 - CH_2$
 $CH_2 - CH_2$

dans laquelle n est un nombre entier de 2 à 4 et Z répond à la formule:

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dans laquelle R₁ et R₂ représentent chacun, indépendamment, un groupe hydrocarboné à chaîne droite ou ramifiée contenant 1 à 34 atomes de carbone de manière que le nombre total d'atomes de carbone dans les groupes R₁ et R₂ soit de 11 à 35, le groupe hydrocarboné Z étant insaturé et formé par réaction de l'acide, de l'anhydride ou d'un ester maléique avec une oléfine interne contenant 12 à 36 atomes de carbone, ladite oléfine interne étant formée par isomérisation de la double liaison oléfinique d'une ou plusieurs alpha-oléfines linéaires de manière à obtenir un mélange d'oléfines internes.

2. Additif soluble dans l'huile, réduisant la friction, suivant la revendication 1, dans lequel R_1 et R_2 sont des groupes hydrocarbonés aliphatiques à chaîne droite ayant au total 15 à 21 atomes de carbone.

3. Additif soluble dans l'huile, réduisant la friction, suivant la revendication 1, qui est le 1-propyltridécénylsuccinimide, le 1-pentyltridécénylsuccinimide, le 1-tridécylpentadécénylsuccinimide ou le 1tétradécyleicosénylsuccinimide.

4. Procédé de préparation d'un additif soluble dans l'huile suivant la revendication 1, procédé qui consiste (a) à isomériser la double liaison oléfinique d'une ou plusieurs alpha-oléfines linéaires contenant 12 à 36 atomes de carbone pour obtenir un mélange d'oléfines internes, (b) à faire réagir le mélange d'oléfines internes avec l'acide, l'anhydride ou un ester maléique pour obtenir un acide, anhydride ou ester succinique à substituant hydrocarboné intermédiaire et (c) à faire réagir le composé intermédiaire avec l'ammoniac pour former un amide, un imide ou un mélange de ces composés.

5. Procédé suivant la revendication 4, dans lequel l'alpha-oléfine consiste principalement en alpha-tétradécène linéaire, en alpha-hexadécène linéaire, en alpha-eicosène linéaire.

6. Procédé suivant la revendication 4 ou la revendication 5, dans lequel l'isomérisation de l'alphaoléfine linéaire est effectuée par chauffage de l'alpha-oléfine linéaire avec un catalyseur acide.

7. Composition d'huile lubrifiante ou de combustible qui contient une quantité réduisant la friction d'un additif suivant l'une quelconque des revendications 1 à 3 ou préparé par un procédé suivant l'une quelconque des revendications 4 à 6.

8. Composition d'huile lubrifiante ou de combustible suivant la revendication 7, dans laquelle l'additif est le tétradécénylsuccinimide, l'hexadécénylsuccinimide, l'octadécénylsuccinimide ou l'eicosénylsuccinimide.

9. Composition d'huile lubrifiante suivant la revendication 7 ou la revendication 8, dans laquelle l'additif est présent en une quantité de 0,05 à 3% en poids.

10. Composition de combustible suivant la revendication 7 ou la revendication 8, dans laquelle l'additif est présent en une quantité de 0,001 à 0,25% en poids.